

DOUBLETIME Plays a Noncatalytic Role To Mediate CLOCK Phosphorylation and Repress CLOCK-Dependent Transcription within the *Drosophila* Circadian Clock[▽]

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Received 20 November 2008/Returned for modification 19 December 2008/Accepted 4 January 2009

Circadian clocks keep time via gene expression feedback loops that are controlled by time-of-day-specific changes in the synthesis, activity, and degradation of transcription factors. Within the *Drosophila melanogaster* circadian clock, DOUBLETIME (DBT) kinase is necessary for the phosphorylation of PERIOD (PER), a transcriptional repressor, and CLOCK (CLK), a transcriptional activator, as CLK-dependent transcription is being repressed. PER- and DBT-containing protein complexes feed back to repress CLK-dependent transcription, but how DBT promotes PER and CLK phosphorylation and how PER and CLK phosphorylation contributes to transcriptional repression have not been defined. Here, we show that DBT catalytic activity is not required for CLK phosphorylation or transcriptional repression and that PER phosphorylation is dispensable for repressing CLK-dependent transcription. These results support a model in which DBT plays a novel noncatalytic role in recruiting additional kinases that phosphorylate CLK, thereby repressing transcription. A similar mechanism likely operates in mammals, given the conserved activities of PER, DBT, and CLK orthologs.

A vast array of animal, plant, and microbial species display daily rhythms of metabolism, physiology, and behavior. These rhythms are controlled by endogenous circadian clocks that are set by ~24-h environmental cycles but persist in the absence of environmental cues. Studies with several phylogenetically diverse model systems have revealed that the circadian time-keeping mechanism is composed of conserved transcriptional feedback loops (reviewed in reference 4). In *Drosophila melanogaster*, these feedback loops are initiated by two basic-helix-loop-helix-PER-ARNT-SIM transcription factors, CLOCK (CLK) and CYCLE (CYC). CLK-CYC heterodimers bind E-box elements during the late day and early evening to activate *period* (*per*) and *timeless* (*tim*) transcription (2, 8, 12, 35). The PERIOD (PER) protein then gradually accumulates during the night as a heterodimer with TIMELESS (TIM) to inhibit CLK-CYC activity (8). The gradual accumulation of PER, an important determinant of circadian period, is controlled by DOUBLETIME (DBT)-dependent phosphorylation (19, 30), which targets PER for degradation in the 26S proteasome (6, 11, 21, 27), and TIM binding, which protects PER from DBT-dependent degradation (11, 19, 21, 30, 31, 34). Once TIM undergoes light- or clock-induced degradation around dawn, PER is degraded and another cycle of CLK-CYC-mediated transcription is initiated.

Although transcriptional repression by PER (or PER-TIM) is essential for feedback loop function, how PER functions to

inhibit CLK-CYC-dependent transcription is not well understood. During the circadian cycle, the accumulation of phosphorylated PER in the nucleus is coincident with hyperphosphorylation of CLK, release of CLK-CYC from E-boxes, and transcriptional repression (41). Given that PER remains bound to DBT when it enters the nucleus and binds CLK (20, 41), it is possible that DBT is also responsible for CLK phosphorylation and transcriptional repression. In support of this possibility, a PER mutant unable to bind DBT (*perΔ*) eliminates both hyperphosphorylation of CLK and repression of CLK-CYC-dependent transcription (17, 29). Nevertheless, *PERΔ* does enter the nucleus and bind CLK, which suggests that PER binding to CLK is not sufficient to remove CLK-CYC from E-boxes and repress transcription (17). These results suggest that DBT is necessary for CLK hyperphosphorylation and transcriptional repression by PER-containing complexes.

If DBT is required for CLK hyperphosphorylation, then CLK should also be hypophosphorylated in the absence of DBT. Although *dbt* null mutants are not viable as adults (19, 30), a *dbt* mutation that severely compromises kinase catalytic activity, *dbt^{ar}*, produces viable adults that lack circadian clock function (33). Surprisingly, CLK is hyperphosphorylated rather than hypophosphorylated in *dbt^{ar}* flies (41). To reconcile the different phosphorylation states of CLK in *perΔ* and *dbt^{ar}* flies, we propose that DBT plays a noncatalytic role in recruiting other kinases into PER repression complexes. Kinases recruited into this complex by DBT phosphorylate CLK, thereby releasing CLK-CYC from E-boxes and repressing transcription.

Here, we demonstrate that entry of DBT into the PER complex is required, but DBT catalytic activity is dispensable for CLK hyperphosphorylation and transcriptional repression. CLK is always hyperphosphorylated when CLK-CYC tran-

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[▽] Published ahead of print on 12 January 2009.

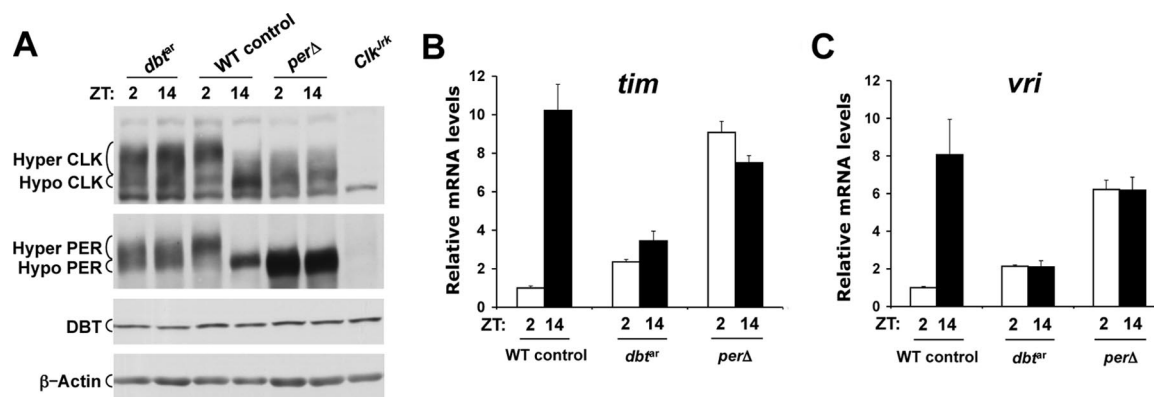


FIG. 1. Comparison of CLK phosphorylation and CLK-CYC transcription levels in *dbt^{ar}* flies with those in *perΔ* flies. (A) Western blot of CLK, PER, DBT, and β -actin levels in *dbt^{ar}/dbt^P* (*dbt^{ar}*), *+/dbt^P* (WT control), *per⁰¹*; *perΔ/dbt^P* (*perΔ*), and *Clk^{rk}* flies collected at ZT 2 or ZT 14. DBT, hypophosphorylated (Hypo) and hyperphosphorylated (Hyper) CLK and PER, and β -actin bands are marked. β -Actin was used as a loading control. (B and C) Quantitative real-time RT-PCR was used to measure *tim* (B) and *vri* (C) mRNA levels in heads from WT control, *dbt^{ar}*, and *perΔ* flies as designated above. The relative mRNA levels were quantified as described in Materials and Methods. Data were plotted as the means \pm standard error of the means ($n = 3$).

scription is repressed, but surprisingly, PER hyperphosphorylation is not required for CLK hyperphosphorylation or repression of CLK-CYC-mediated transcription. These results strongly support a novel noncatalytic role for DBT in recruiting other kinases that phosphorylate CLK and promote transcriptional repression, indicating that PER phosphorylation by DBT is not a prerequisite for transcriptional repression.

MATERIALS AND METHODS

Fly stocks. The *w¹¹¹⁸* strain served as a wild-type (WT) control for clock function. The UAS-*dbt^{K/R}* (*dbt^{K/R}*) is the *dbt* mutant lacking kinase catalytic activity due to a K38R substitution, *perΔ* (*per⁰¹*, *w¹¹¹⁸*; *perΔ-HAHIS/+*), and *tim-Gal4* transgenic strains were described previously (10, 17, 26). The UAS-*dbt^{K/R}* line UAS-KR22, the *perΔ* line *perΔF21*, and the *tim-Gal4* line 62 were used. The *dbt^{ar}* mutant employed in this study has the genotype *dbt^{ar}/dbt^P*, where *dbt^P* is a homozygous lethal P element insert that is apparently *dbt* null (19, 30). We used *dbt^{ar}/dbt^P* flies because they are healthier than homozygous *dbt^{ar}* flies in our hands and they have the lowest level of DBT catalytic activity as viable adults. *dbt^{ar}/dbt^P* flies were produced by crossing *dbt^{ar}/TM3* flies to *dbt^P/TM3* flies. The *perΔ* genotype employed here (*per⁰¹*, *w¹¹¹⁸*; *perΔ/dbt^P*) was generated to keep the genetic background comparable to that of the *dbt^{ar}* genotype (*dbt^{ar}/dbt^P*) by crossing *per⁰¹*, *w¹¹¹⁸*; *dbt^P/TM3* flies with *per⁰¹*, *w¹¹¹⁸*; *perΔ/MKRS* flies. The *per⁰¹*, *w¹¹¹⁸*; *perΔ*, *dbt^{ar}/dbt^P* double mutants, referred to as *perΔ*, *dbt^{ar}* in the text, were produced by recombining the *perΔ* transgene onto a third chromosome containing *dbt^{ar}*. Recombinant third chromosomes bearing the *perΔ* insert were identified as *dbt^{ar}* by their loss of a FokI restriction site caused by the *dbt^{ar}* mutation and were verified by DNA sequencing. The *per⁰¹*, *w¹¹¹⁸*; *perΔ*, *dbt^{ar}/TM6B* flies were crossed to *per⁰¹*, *w¹¹¹⁸*; *dbt^P/TM3* flies to generate *per⁰¹*, *w¹¹¹⁸*; *perΔ*, *dbt^{ar}/dbt^P* double-mutant flies. UAS-*dbt^{K/R}/tim-Gal4*; *dbt^{ar}/dbt^P* flies, referred to as *dbt^{K/R}*; *dbt^{ar}* flies in the text, were generated by crossing *dbt^{K/R}/CyO*; *dbt^{ar}/dbt^P* flies with *tim-Gal4/CyO*; *dbt^{ar}/dbt^P* flies. The *dbt^{K/R}/CyO*; *dbt^{ar}/dbt^P* and *tim-Gal4/CyO*; *dbt^{ar}/dbt^P* flies were used as *dbt^{ar}* no-driver and *dbt^{ar}* no-responder controls, respectively.

Western blotting. For preparing fly head extract, flies were entrained in a 12-h light/12-h dark (LD) incubator for at least 3 days and collected at the indicated time points. Isolated frozen fly heads were homogenized in radioimmunoprecipitation assay (RIPA) buffer (20 mM Tris at pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.05 mM EGTA, 10% glycerol, 1% Triton X-100, 0.4% sodium deoxycholate, 0.1% SDS [sodium dodecyl sulfate]) containing 0.5 mM PMSF (phenylmethylsulfonyl fluoride), 10 μ g/ml aprotinin, 10 μ g/ml leupeptin, 2 μ g/ml pepstatin A, 1 mM Na_3VO_4 , and 1 mM NaF (hereafter referred to as the protease inhibitor mixture). This homogenate was sonicated 5 to 10 times for 10 s each time, using a Misonix XL2000 model sonicator at a setting of 3 and then centrifuged at $20,000 \times g$ for 10 min. The supernatant was collected as RIPA S extract, and protein concentration was determined by the Bradford assay. Equal

amounts of RIPA S extract were run, transferred, and probed with antibodies as follows: guinea pig anti-CLK (GP-47), 1:2,000 (15); guinea pig anti-PER (GP-73), 1:3,000 (37); rat anti-TIM (TR3), 1:2,000 (37); rabbit anti-DBT, 1:4,000 (26); and mouse anti- β -actin (Abcam), 1:2,000. Horseradish peroxidase-conjugated secondary antibodies (Sigma) against guinea pig, rat, rabbit, and mouse were diluted 1:1,000. Immunoblots were visualized using ECL plus (Amersham) reagent.

Quantitative real-time RT-PCR. Total RNA was isolated from frozen fly heads, using Trizol (Invitrogen), and treated with a Turbo DNase DNA-free kit (Ambion) to eliminate genomic DNA contamination. DNA-free total RNA (1.0 μ g) was reverse transcribed using oligo(dT)₁₂₋₂₈ primers (Invitrogen) and Superscript II (Invitrogen). The reverse transcription (RT) product was amplified and analyzed with an Applied Biosystems model 7500 Fast real-time PCR system using Power SYBR green PCR Master Mix (Applied Biosystems) and gene-specific primers. All primer pairs were designed to span an exon-intron boundary to prevent genomic DNA amplification. The gene-specific primer pairs used for *rp49* were 5'-TACAGGCCCAAGATCGTGAA-3' and 5'-GCACTCTGTGTGTCGATACCC-3'; for *vri* were 5'-ATGAACAACGTCGGCTATC-3' and 5'-CTGCGGACTTATGGATCCTC-3'; for *per* were 5'-TGATGGGCGACTACAAC TCC-3' and 5'-GTCGCTATTCCTCATTTGCTGT-3'; and for *tim* were 5'-GGTG GCATCTGTGTACGAAA-3' and 5'-GATCTCGGTCGCTCAAGTC-3'. For each sample, RNA quantity was determined by the standard curve for each gene that was analyzed. The RNA quantity of *vri*, *per*, or *tim* was normalized to that of *rp49*. For each data series, the *rp49*-normalized values were further normalized to levels in *dbt^P/+* (Fig. 1) and *w¹¹¹⁸* (see Fig. 4) at zeitgeber time 2 (ZT 2) (during LD cycles, lights on is referred to as ZT 0 and lights off is referred to as ZT 12) to yield relative mRNA levels.

IP. For immunoprecipitation (IP) assays, EB3-S extract was prepared by homogenizing frozen fly heads in EB3 buffer (10 mM HEPES at pH 7.5, 5 mM Tris at pH 7.5, 50 mM KCl, 1 mM EDTA, 0.05 mM EGTA, 10% glycerol, and 0.1% Triton X-100) containing protease inhibitor mixture, sonicated as described for preparing RIPA S extracts, and centrifuged at $20,000 \times g$ for 10 min, and the supernatant was collected as EB3-S extract. To 1 mg of the EB3-S extract, 2.5 μ l of DBT antiserum was added, and the mixture was incubated at 4°C overnight. Immune complexes were collected by incubating 35 μ l of 50% protein A-Sepharose beads (Amersham) at 4°C for 2 h and then washing the beads three times with EB3 buffer. The immunoprecipitates were eluted by boiling in 15 μ l of 2 \times loading buffer and then run in parallel with 50 μ g of EB3-S extracts as input. Immunoblots were processed as described previously for Western blotting.

ChIP. Chromatin IP (ChIP) assays were performed as described previously (41), with modifications. Briefly, frozen fly heads were homogenized and cross-linked in ~ 5 volumes of HX buffer (50 mM HEPES at pH 8.0, 140 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 0.4% Igel CA-630, 0.2% Triton X-100, 1% formaldehyde, 1 mM PMSF, 1 mM Na_3VO_4 , and 1 mM NaF) for 10 min at 25°C. Glycine was added to a final concentration of 0.125 M to stop the cross-linking reaction. The homogenates were then filtered with 100- μ m nylon mesh. The nuclei were harvested by centrifugation at $800 \times g$ for 5 min and washed three

times with XN wash buffer (20 mM Tris at pH 7.5, 150 mM NaCl, 1 mM EDTA, and 0.5 mM EGTA) containing protease inhibitor mixture (see “Western blotting” above). The cross-linked nuclei were suspended in RIPA buffer containing protease inhibitor mixture and sonicated 15 times for 10 s each time using a Misonix model XL2000 sonicator at a setting of 3. After centrifugation at $25,000 \times g$ for 10 min, the supernatant was collected as XN extract and quantified by the Bradford assay. An aliquot of 50 μ g of XN extract was stored at -80°C as input. For IP, 500 μ g of XN extract was diluted in 4 volumes of IP buffer (20 mM Tris at pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 1% Triton X-100, and 0.01% SDS) containing protease inhibitor mixture. The diluted XN extract was precleared as described previously (41) and then immunoreacted overnight by the addition of 3 μ l of anti-PER (GP-73) and salmon sperm DNA to a final concentration of 0.1 μ g/ μ l. Immune complexes were recovered, and DNA was purified as described previously (41).

Quantitative real-time PCR was used to measure E-box-containing DNA fragments. The suspended DNA samples were further diluted at 1:40 for ChIP DNA and at 1:600 for input DNA for real-time PCR. The circadian regulatory sequence (CRS) containing a fragment upstream of *per* and an E-box-containing fragment upstream of *cry* were amplified and quantified as described above in “Quantitative real-time RT-PCR” using the primer pair 5'-CCAGTGCCAGTGCAGATTC-3' and 5'-GATGCCAAGTGTCAATCCAAGC-3' for the CRS E-box; and the primer pair 5'-CCCCTTATAATCTGGTTTGG-3' and 5'-TGCTAGACAACGACAACGAC-3' for the *cry* E-box. The *cry* E-box is not bound by CLK-CYC (data not shown) and was used as a measure of background PER binding. The percentage of total DNA was calculated by the following formula: the percentage of total DNA = (CRS quantity from ChIP \times 40)/(CRS quantity from input \times 10 \times 600) \times 100% - (cry E-box quantity from ChIP \times 40)/(cry E-box quantity from input \times 10 \times 600) \times 100%.

RESULTS

PER-DBT binding, but not DBT catalytic activity, is required for CLK hyperphosphorylation and transcriptional repression. We first confirmed PER and CLK phosphorylation levels in *per* Δ and *dbt*^{ar} flies (see Materials and Methods for complete genotypes). The small deletion within the PER Δ protein prohibits DBT binding and eliminates PER and CLK phosphorylation and transcriptional repression, even though PER Δ is able to bind CLK and TIM (16, 17). In contrast, catalytically compromised DBT^{ar} protein shows high levels of PER and CLK phosphorylation, though its effect on transcription was not tested (33, 41). As expected, phosphorylation of PER and CLK is strongly skewed toward hyperphosphorylated forms in *dbt*^{ar} flies and is almost entirely hypophosphorylated in *per* Δ flies (Fig. 1A). Moreover, although DBT^{ar} is catalytically compromised (18, 33), it accumulates to levels that are similar to those of WT DBT (Fig. 1A), as previously seen in *tim*⁰¹; *dbt*^{ar} flies (7). Since PER and CLK are hypophosphorylated when CLK-CYC-mediated transcription is activated and are hyperphosphorylated when CLK-CYC-mediated transcription is repressed (41), we determined the *tim* and *vri* mRNA levels in *per* Δ and *dbt*^{ar} flies. Both *tim* and *vri* mRNA levels are high in *per* Δ flies (Fig. 1B and C), consistent with increased CLK-CYC-mediated transcription of *per* when PER and CLK are hypophosphorylated in WT and *per* Δ flies (17, 41). In contrast, *tim* and *vri* mRNAs were at low levels in *dbt*^{ar} flies (Fig. 1B and C), thereby strengthening the link between PER and CLK hyperphosphorylation and repression of CLK-CYC-mediated transcription. A similar decrease in *tim*, *vri*, and *per* mRNA levels was detected in *tim*⁰¹; *dbt*^{ar} flies (7) and was likely due to severely reduced DBT catalytic activity.

In WT flies, PER-DBT complexes bind to CLK-CYC, promote CLK hyperphosphorylation, and release CLK-CYC from E-boxes (23, 41). In *per* Δ mutants, PER Δ forms a complex with CLK-CYC but it neither promotes CLK hyperphosphorylation

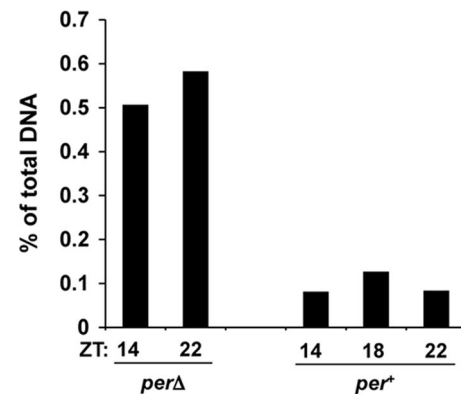


FIG. 2. DBT is required for PER to remove CLK-CYC from E-boxes. ChIP assays were carried out to probe PER binding to the *per* CRS E-box in *per* Δ and *per*⁺ control flies at the indicated times, using PER (GP-73) antiserum. The percentage of total CRS E-box DNA bound by PER was calculated as described in Materials and Methods.

nor removes CLK-CYC from E-boxes (17). Consequently, PER Δ -CLK-CYC complexes are expected to remain associated with E-boxes. ChIP assays using PER antiserum show that PER is always associated with the *per* CRS E-box (12, 13) in *per* Δ flies, but not in *per*⁺-rescued control flies (Fig. 2). ChIP experiments using antihemagglutinin (anti-HA) antibody to probe PER-HA interactions with the *per* CRS E-box in *per* Δ and *per*⁺-rescued control flies produced similar results (data not shown). Thus, even though PER is bound to CLK-CYC in *per* Δ flies, CLK is not phosphorylated and CLK-CYC is not released from E-boxes if DBT cannot enter the PER-CLK-CYC complex. Although PER levels in *per* Δ flies are much higher than those in WT flies, PER levels in WT flies are already saturated with respect to CLK (3), which argues that PER interactions with the CRS E-box are not due simply to high PER levels in *per* Δ flies. The inability of PER to immunoprecipitate E-box complexes in WT flies suggests that PER binding rapidly removes CLK-CYC from the E-box.

DBT^{ar} binds PER to promote CLK phosphorylation. We previously showed that CLK hyperphosphorylation is PER dependent (41). If DBT binds PER to recruit other kinases that mediate CLK hyperphosphorylation, DBT^{ar}-dependent hyperphosphorylation of CLK should also require PER-DBT^{ar} binding. Alternatively, mutant DBT^{ar} protein may bypass the requirement for PER-DBT binding to mediate CLK hyperphosphorylation. To determine whether hyperphosphorylation of CLK requires PER-DBT^{ar} binding, we generated *per* Δ , *dbt*^{ar} double mutants (described in Materials and Methods) and examined PER and CLK phosphorylation. PER and CLK remain hypophosphorylated in *per* Δ , *dbt*^{ar} flies, which phenocopies the *per* Δ rather than the *dbt*^{ar} mutant (Fig. 3). These results indicate that DBT^{ar} binds PER to bring about PER and CLK hyperphosphorylation and thus support a noncatalytic role for DBT.

The dependence of PER and CLK hyperphosphorylation on PER-DBT^{ar} binding implies that DBT^{ar} is present in a complex with PER and CLK-CYC. To determine if this is the case, DBT antiserum was used to immunoprecipitate head extracts from *dbt*^{ar}, WT, and *per* Δ flies collected at different times of day. DBT coimmunoprecipitated PER and CLK only when

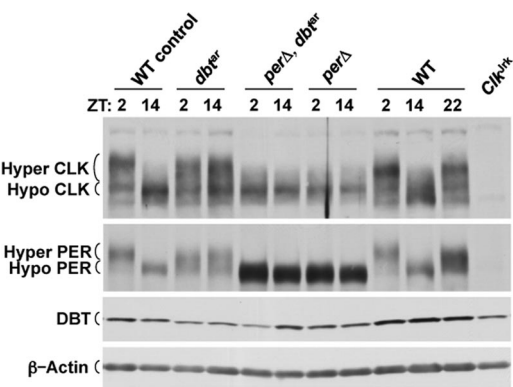


FIG. 3. DBT^{ar} promotes CLK hyperphosphorylation (Hyper) via interactions with PER complexes. Western blot shows CLK, PER, DBT, and β -actin levels in the heads of +/*dbt*^P (WT control), *dbt*^{ar}/*dbt*^P (*dbt*^{ar}), *per*⁰¹;; *per* Δ , *dbt*^{ar}/*dbt*^P (*per* Δ , *dbt*^{ar}), *per*⁰¹;; *per* Δ /*dbt*^P (*per* Δ), *w*¹¹¹⁸ (WT), and *Clk*^{Jrk} flies collected at ZT 2 or ZT 14. DBT, hypophosphorylated (Hypo) and hyperphosphorylated (Hyper) CLK and PER, and β -actin bands are marked. β -Actin was used as a loading control.

PER and CLK were hyperphosphorylated, as determined at ZT 2 and ZT 14 in *dbt*^{ar} flies and at ZT 22 and ZT 2 in WT flies (Fig. 4). However, DBT did not coimmunoprecipitate PER and CLK when PER and CLK were hypophosphorylated at ZT 2 and ZT 14 in *per* Δ flies and at ZT 14 in WT flies (Fig. 4). These data demonstrate that DBT^{ar} is indeed in a complex with PER and CLK in *dbt*^{ar} flies and confirm previous results showing that PER-DBT-CLK-CYC complexes form in phase with PER and CLK hyperphosphorylation and transcriptional repression in WT flies but lack DBT in *per* Δ flies (17, 41). Taken together with the PER-DBT^{ar} binding required for DBT^{ar}-dependent PER and CLK hyperphosphorylation (Fig. 3), these results suggest that DBT recruits or DBT and PER jointly recruit other kinases that phosphorylate PER and CLK to repress transcription.

Blocking DBT catalytic activity in clock cells prevents PER phosphorylation but not CLK hyperphosphorylation and CLK-CYC transcriptional repression. Although results from *dbt*^{ar} flies strongly support a noncatalytic role for DBT in PER

and CLK phosphorylation and transcriptional repression, it is possible that DBT^{ar} retains a modified kinase activity that is still capable of phosphorylating PER and CLK with altered specificity (33), consistent with recent results showing low levels of casein phosphorylation by DBT^{ar} (18). Phosphorylation of different target sites can alter protein activity, localization, and/or degradation. For instance, phosphorylation of Ser 659 on mPER2 promotes mPER2 stabilization, whereas phosphorylation of other sites by CK1 δ/ϵ promotes mPER2 degradation (40). Likewise, DBT phosphorylates S47 on PER to promote degradation (6), whereas other PER phosphorylation sites are predicted to control subcellular localization (1, 25, 38). To determine whether PER and CLK phosphorylation in *dbt*^{ar} flies is due to other kinases or to altered DBT^{ar} target site specificity, we employed a dominant negative form of DBT (DBT^{K38R}) that lacks kinase catalytic activity due to a K38R substitution (26). Since *dbt* null mutants die during development (19, 30), DBT^{K38R} (hereafter referred to as DBT^{K/R}) is expressed in a tissue-specific manner using the Gal4/UAS system, thereby avoiding tissues that give rise to developmental lethality. Overexpressing DBT^{K/R} in vitro and in oscillator cells from WT flies antagonizes PER hyperphosphorylation, thus demonstrating that DBT mediates the vast majority of PER phosphorylation (26). PER is hypophosphorylated in flies that overexpress DBT^{K/R} in oscillator cells (26), which contrasts with PER hyperphosphorylation in *dbt*^{ar} flies (Fig. 1 and 3). This difference in PER phosphorylation implies that DBT^{ar} retains reduced and/or off-target catalytic activity that phosphorylates PER.

To produce the greatest reduction in DBT catalytic activity, *tim*-Gal4 and UAS-*dbt*^{K/R} transgenes were used to overexpress DBT^{K/R} in oscillator cells from *dbt*^{ar} flies (referred to as *dbt*^{K/R}; *dbt*^{ar} flies). In *dbt*^{K/R}; *dbt*^{ar} flies, the high levels of DBT^{K/R} compared to that of DBT^{ar} suggest that DBT^{K/R} will effectively block DBT^{ar} catalytic activity (Fig. 5A). Indeed, PER levels are constitutively high and are almost completely hypophosphorylated in *dbt*^{K/R}; *dbt*^{ar} flies (Fig. 5A, compare lanes 3 and 4 to lanes 1 and 2, 5 and 6, or 11 and 12). In contrast, CLK is constitutively hyperphosphorylated in *dbt*^{K/R}; *dbt*^{ar} flies (Fig. 5A, compare lanes 3 and 4 to lanes 1 and 2, 5

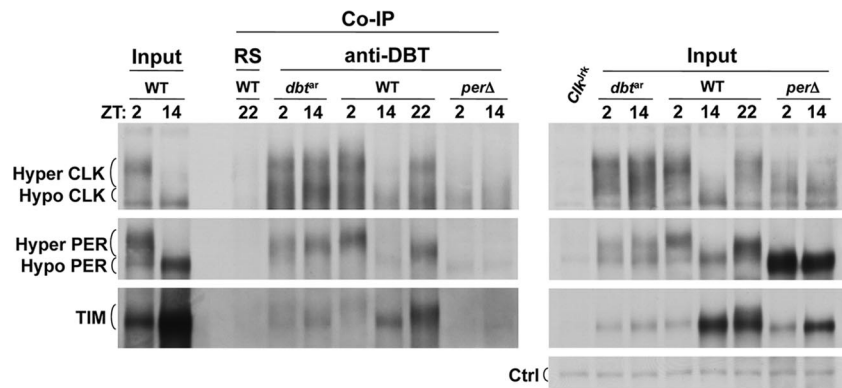


FIG. 4. DBT is in a complex with PER, TIM, and CLK in *dbt*^{ar} but not *per* Δ flies. EB3-S extracts from *w*¹¹¹⁸ (WT), *dbt*^{ar}/*dbt*^P (*dbt*^{ar}), and *per*⁰¹;; *per* Δ /+ (*per* Δ) fly heads collected at the indicated time points were subjected to coimmunoprecipitations (Co-IP) using a rabbit DBT antiserum (anti-DBT) or normal rabbit serum (RS). Western blots containing immunoprecipitates and input extracts (Input) were probed with CLK, PER, and TIM antibodies, respectively. DBT and hypophosphorylated (Hypo) and hyperphosphorylated (Hyper) CLK and PER bands are marked. Ctrl is a constant nonspecific band detected by PER antiserum as a loading control.

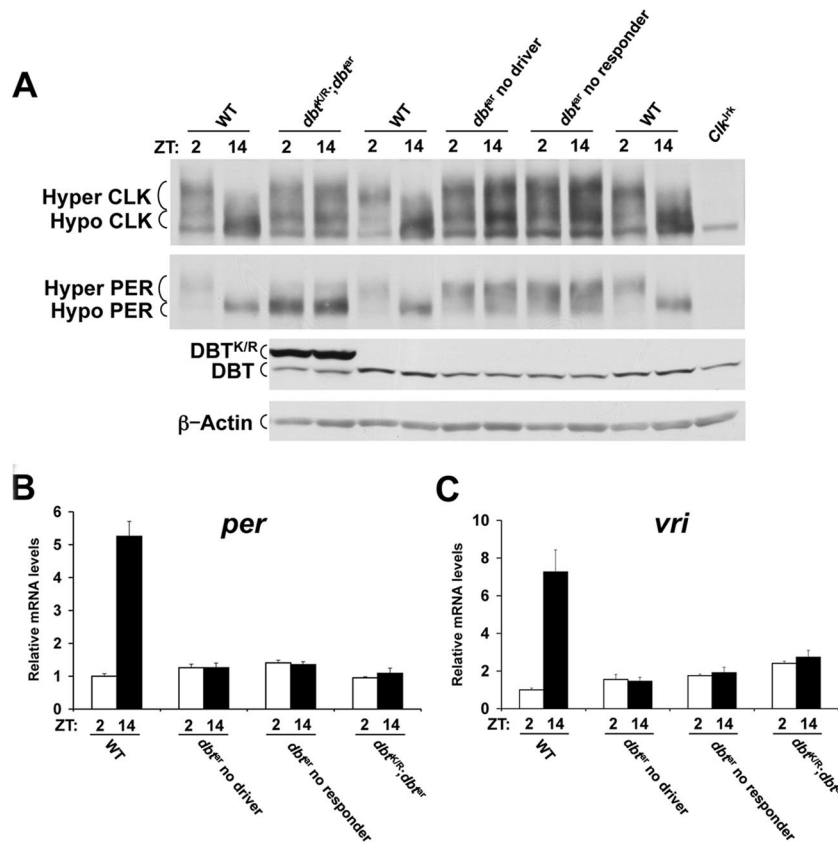


FIG. 5. Blocking DBT catalytic activity in oscillator cells prevents PER phosphorylation but not CLK hyperphosphorylation and CLK-CYC transcriptional repression. (A) Western blot of CLK, PER, DBT, and β -actin levels in *w¹¹¹⁸* (WT), *UAS-dbt^{K/R}/tim-Gal4; dbt^{ar}/dbt^P* (*dbt^{K/R}; dbt^{ar}*), *UAS-dbt^{K/R}/+*; *dbt^{ar}/dbt^P* (*dbt^{ar} no driver*), *tim-Gal4/+; dbt^{ar}/dbt^P* (*dbt^{ar} no responder*), and *Clk^{trk}* flies collected at the indicated times. DBT, DBT^{K/R}, hypophosphorylated (Hypo) and hyperphosphorylated (Hyper) CLK and PER, and β -actin bands are marked. β -Actin was used as a loading control. (B and C) Quantitative real-time RT-PCR was used to measure *per* (B) and *vri* (C) mRNA levels in heads from WT, *dbt^{K/R}; dbt^{ar}*, *dbt^{ar} no driver*, and *dbt^{ar} no responder* flies as designated above. The relative mRNA levels were quantified as described in Materials and Methods. The data were plotted as the means \pm standard error of the means ($n = 3$).

and 6, or 11 and 12), which further supports our argument that DBT catalytic activity is not necessary for CLK hyperphosphorylation. Given that PER hypophosphorylation is associated with CLK-CYC transcriptional activation and that CLK hyperphosphorylation is associated with CLK-CYC transcriptional repression, we wanted to determine the level of CLK-CYC-dependent transcription in *dbt^{K/R}; dbt^{ar}* flies. The levels of *per* and *vri* mRNA are near that of the WT trough in *dbt^{K/R}; dbt^{ar}* fly heads (Fig. 5B and C), thus reinforcing the link between CLK hyperphosphorylation and repression of CLK-CYC-dependent transcription and severing the association between PER hypophosphorylation and activation of CLK-CYC-dependent transcription.

DISCUSSION

Nonphosphorylated PER accumulates to high levels in homozygous *dbt^P* null mutant larvae and *dbt^{K/R}* adults due to the lack of DBT catalytic activity (26, 30). Despite the high levels of PER in *dbt^P* larvae, CLK-CYC-dependent transcription is not repressed by monomeric PER in LD cycles (30), consistent with results for *per Δ* flies and the requirement that DBT recruit additional kinases that phosphorylate CLK. The

high levels of nonphosphorylated PER that accumulate in *dbt^{K/R}; dbt^{ar}* flies indicate that DBT catalytic activity has been eliminated. Nevertheless, CLK is hyperphosphorylated in *dbt^{K/R}; dbt^{ar}* flies, thus demonstrating that DBT catalytic activity is dispensable for CLK hyperphosphorylation. Conversely, DBT catalytic activity is required for PER hyperphosphorylation. The presence of phosphorylated PER in *dbt^{ar}* flies implies that DBT^{ar} retains catalytic activity but that this activity is defective since it leads only to the partial degradation of PER (Fig. 1 and 3 to 5). The remaining phosphorylated PER apparently forms a PER repression complex sufficient to sustain transcriptional repression.

The CLK phosphorylation state parallels transcriptional activity in the WT, *per* mutant, and *dbt* mutant genotypes; CLK is hyperphosphorylated when CLK-CYC-dependent transcription is repressed and is hypophosphorylated when CLK-CYC-dependent transcription is activated (41) (Fig. 1 and 5). In contrast, results with *dbt^{K/R}; dbt^{ar}* flies demonstrate that PER hyperphosphorylation is not a prerequisite for transcriptional repression and strengthen the argument that DBT plays a noncatalytic role in PER complexes to mediate transcriptional repression. The loss of CLK-CYC activity in *dbt^{K/R}; dbt^{ar}* flies suggests that CLK is phosphorylated at the same sites that

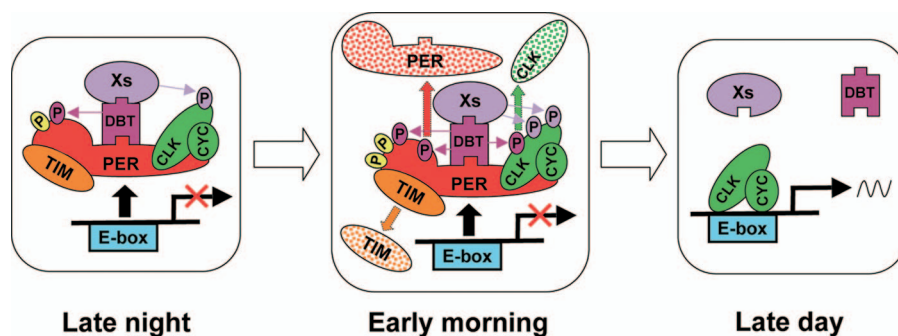


FIG. 6. Model of noncatalytic DBT function during the circadian cycle (see text for description). Transcriptional regulatory events occurring during late night (left panel), early morning (middle panel), and late day (right panel) are shown. PER, red shape; TIM, orange oval; DBT, plum shape; CLK-CYC, green ovals; DBT-bridged CLK kinases, Xs in purple oval; DBT phosphorylation, plum-colored oval with a P; DBT-bridged CLK kinase phosphorylation, purple oval with a P; PER phosphorylation by other kinases, yellow oval with a P; E-box regulatory element, blue rectangle; transcriptional repression, arrow with red X; transcriptional activation, arrow with black waveform; PER degradation, red arrow to red stippled PER; TIM degradation, orange arrow to orange stippled TIM; CLK degradation, green arrow to green stippled CLK; inhibition of CLK-CYC binding to E-boxes, black arrow.

repress transcriptional activity in WT flies, thus implying that the same kinases mediate CLK phosphorylation in the WT and in the *dbt^{K/R}*; *dbt^{ar}* mutant strains. Furthermore, the 270-amino-acid CLK-CYC interaction domain of PER, which contains the DBT binding region (17, 29), is sufficient to inhibit CLK-CYC transcription in cell culture (5), consistent with a requirement for DBT, but not PER phosphorylation, in repressing CLK-CYC transcription. Likewise, eliminating DBT expression via RNA interference in cell culture gives rise to high levels of nonphosphorylated PER that are unable to repress CLK-CYC-dependent transcription (28), further supporting a requirement for DBT in repressing CLK-CYC-mediated transcription.

Our results demonstrate that DBT catalytic activity is not necessary for phosphorylating CLK or repressing CLK-CYC-mediated transcription, but how DBT carries out these activities is not known. We speculate that DBT forms a physical bridge to recruit other kinases that phosphorylate CLK and repress CLK-CYC-dependent transcription (Fig. 6). During the early evening, PER destabilization by DBT phosphorylation and stabilization by TIM binding result in the gradual accumulation of PER (11, 19, 21, 30, 31, 34). PER phosphorylation by CK2 then promotes the nuclear localization of PER-DBT and PER-DBT-TIM complexes (1, 25, 38), where continued phosphorylation by DBT produces hyperphosphorylated PER (6, 30). As PER-DBT-TIM complexes accumulate in the nucleus late at night, they bind CLK-CYC via interactions between PER and CLK (3, 22, 23). Once the PER-DBT-TIM-CLK-CYC complex is formed, DBT recruits other kinases that hyperphosphorylate CLK, thereby releasing CLK-CYC from E-boxes and repressing transcription (39, 41). Although DBT may recruit other kinases via direct binding (Fig. 6), it is also possible that entry of DBT into the PER repression complex could promote binding of other kinases to PER and/or CLK, which then directly phosphorylate CLK. CLK phosphorylation coincides with transcriptional repression, which implies that CLK phosphorylation represses transcription. However, experimental support for this possibility awaits identification of CLK phosphorylation sites and/or the kinases that phosphorylate CLK. After dawn, light or clock-dependent

degradation of TIM permits SLIMB binding of phosphorylated PER S47 (6), which targets PER for degradation in the 26S proteasome (11, 21, 27). Phosphorylation of CLK by DBT also triggers CLK degradation (16, 41), and by mid day, hypophosphorylated CLK derived from new synthesis and/or dephosphorylation binds E-boxes to initiate the next cycle of transcription.

Loss of DBT catalytic and noncatalytic activities together, or DBT catalytic activity alone, stops the oscillator at different circadian phases. The loss of PER-DBT binding in flies expressing *PERΔ* disables both DBT catalytic and bridging functions, thus stopping the feedback loop during the early evening when CLK is hypophosphorylated and CLK-CYC-mediated transcription is high. In contrast, loss of DBT catalytic function alone in *dbt^{ar}* and *dbt^{K/R}*; *dbt^{ar}* flies stops the feedback loop during the early morning, when CLK is hyperphosphorylated and CLK-CYC-mediated transcription is low. The different phases of oscillators stopped by the loss of DBT catalytic or catalytic plus noncatalytic functions suggest that DBT catalytic activity is required for progression from transcriptional repression to activation.

The DBT bridge model may be applicable to eukaryotic clocks in general. Like CLK in *Drosophila*, the feedback loop activators WHITE COLLAR 1 and WHITE COLLAR 2 (WCC) in *Neurospora* and CLOCK (as a heterodimer with BMAL1) in mammals are transcriptionally active when they are hypophosphorylated and repressed when they are hyperphosphorylated (24, 32, 36). CK1 and CK2 mediate FREQUENCY (FRQ)-dependent hyperphosphorylation of the WCC in *Neurospora Crassa* to repress transcription (14), but whether these processes are mediated by catalytic activity or by bridging is not known. In mammals, CRYPTOCHROME-PERIOD (CRY-PER) complexes contain CK1ε and CK1δ and promote CLOCK hyperphosphorylation and transcriptional repression (24). As in *Drosophila*, expression of dominant negative CK1ε represses CLOCK-BMAL1-dependent transcription in cultured mammalian cells (9). Although the state of CLOCK phosphorylation in these cells was not determined, the strong link between CLOCK phosphorylation and transcriptional repression argues for a CK1ε bridging function.

Additional experiments will be necessary to support or rule out a role for CK1 bridging in *Neurospora* and mammals.

ACKNOWLEDGMENTS

We thank Isaac Edery for providing PER and TIM antisera. We also thank members of the Hardin laboratory for helpful discussions.

This work was supported by NIH grants NS051280 (P.E.H.) and MH056895 (J.L.P.).

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